PATENT SPECIFICATION

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Inventor: DONALD MOORE JONAS

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COMPLETE SPECIFICATION

Synthetic Thermoplastic Polymer Composition

We, BERK LIMITED, a British Company of Berk House, 8 Baker Street, London W.1., do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement: -

This invention is concerned with synthetic thermoplastic polymer compositions which contain flame-retardants and have drip-resistant

properties.

The synthetic thermoplastic polymers with which this invention is concerned (hereinafter referred to, for convenience, simply as "syn-15 thetic thermoplastic polymers") are polyamides, polystyrene and polyolefins derived from straight chain olefins. These polymers are combustible and, in addition to being combustible, melt and drip readily when heated beyond their softening point. While the flammability of these polymers can be substantially reduced by the incorporation therein of one or more flame retardants, they represent (in the absence of appropriate further additives) a considerable fire hazard even when they contain flame retardants since when heated beyond their softening point, they will melt and hot material will drip or flow therefrom onto surrounding areas and increase the risk of ignition of surrounding materials even though the polymer itself may not burn. Examples of such polymers are, in addition to polystyrene, polyethylene, polypropylene and nylon. We have now found that the tendency of

these synthetic thermoplastic polymers to melt and drip when heated to temperatures above their softening points can be substantially reduced by incorporating therein a small proportion (that is less than 10%, based on the 40 weight of the polymer) of an organophilic cation modified clay. The present invention accordingly comprises a polymer composition comprising a synthetic thermoplastic polymer (as herein defined) and, incorporated therein, one or more flame retardants and up to 10%, based on the weight of the polymer, of an

organophilic cation modified clay.

Suitable organophilic cation modified clays for use in accordance with the invention are cationic clays of medium or high cation ex- 50 change capacity, the cation of which is replaced by a so-called onium base. The onium base may contain a pentavalent atom as in the cases of ammonium, phosphonium, arsonium and stibonium bases; a tetravalent atom as in 55 the cases of oxonium, sulphonium, selenonium, telluronium and stannonium bases; or a trivalent atom as in the case of an iodonium base. The cation exchange capacity is generally reported as the number of milliequivalents of 60 exchangeable base which can be exchanged per 100 grams of clay dried at 105° C. The cationic clays have very different cation-exchange capacities, the numerical values varying from about 3 to about 120 depending upon the type of clay. On this basis clays have been empirically divided into those of low cationexchange capacity, i.e. up to about 15; those of medium cation-exchange capacity, i.e. above 15 but below 40; and those of high cationexchange capacity, i.e. 40 and above.

Examples of clays having a medium cationexchange capacity are sepiolite and attapulgite. Examples of clays having a high cationexchange capacity are the montmorillonites, such as sodium, potassium, lithium and other bentonites, particularly those of the Wyoming and South Dakota (U.S.A.) types and magnesium bentonite (which is also known as hectorite), saponite and nontronite. Certain socalled "synthetic clays" are also known and provided that they are equivalent to the natural clays described above in their ability to react with onium compounds, they are suitable.

Details of the preparation of organophilic 85 cation modified clays are given, for instance,

in British Specification No. 664,830 and U.S. Patent No. 2,531,440. If desired, the sodium ion content of the clay may be increased prior to reaction with the onlum compound as described in British Specification No. 904,880.

Specific examples of suitable organophilic cation modified clays are: dimethyl dioctadecyl ammonium sepiolite, octadecyl ammonium montmorillonite, dimethyl dioctadecyl ammonium montmorillonite, and dimethyl di-

octadecyl ammonium hectorite.

Another class of organophilic cation modified clays that can be used are cationic clays of high cation-exchange capacity, e.g. montmorillonites, which are associated with an aliphatic amine containing at least 12 carbon atoms, such as dodecylamine, octadecylamine, methyloctadecylamine, dioctadecylamine and dimethyloctadecylamine, or a derivative thereof, for example the derivatives of such amines obtained by fusing them with a non-ionic, long chain, organic polar compound, such as stearamide, stearonitrile and cetyl alcohol.

A wide variety of flame retardants, both organic and inorganic and mixtures thereof, can be used in the polymer compositions. The flame retardants can be halogen-containing, examples of flame retardants of this type being the chlorinated parathus sold under the trade 30 mark "Cereclor", tris-β-chloroethyl phosphate, tribromophenol, ammonium chloride and ammonium bromide (the ammonium halides may be coated with an organophilic coating, such as stearic acid, prior to incorporation in the polymer composition). Suitable halogen-free flame retardants are, for example, phosphoric acid esters, such as tri-tolyl phosphate, antimony trioxide and arsenic trioxide.

The proportion of total flame retardant may suitably range from 0.5 up to 50%, based on the weight of the polymer. The proportion of flame retardant used will depend for instance upon the nature of the polymer into which it is to be incorporated and the degree of flame retardancy required in the polymer composition. The proportion of organophilic cation modified clay is suitably from 0.5 to 5%, based on the weight of the polymer, although higher proportions may be desirable in some cases.

The polymer compositions can also, of course, contain one or more plasticisers, antioxidants, stabilisers, anti-static agents, fillers, pigments and the like, but the amounts of such additional materials should not be such as to detract deleteriously from the flame-retardant and drip resistant character of the composition as a whole.

The organophilic cation modified clay can be added as a dry powder directly to the unplasticised synthetic thermoplastic polymer during processing of the polymer. If desired, the organophilic cation modified clay may be dispersed in a volatile liquid organic vehicle prior to incorporation in the polymer; examples of such vehicles include trichloroethylene,

toluene, xylene and white spirit A masterbatch of polymer and organophilic cation modified clay may be used as the method of introducing the organophilic cation modified clay; such a masterbatch may be made by dispersing the organophilic cation modified clay in a volatile liquid organic vehicle, such as toluene, and then incorporating the dispersion in a proportion of polymer which may itself be dispersed in an organic liquid vehicle. The volatile organic liquid vehicle may be substantially completely removed prior to or during processing of the unplasticised synthetic thermoplastic polymer. The organophilic cation modified clay may, on the other hand, be dispersed in a relatively non-volatile vehicle, including, for instance, the additives for the polymer, such as the flame retardant, andvehicles such as paraffin wax and a low molecular weight polyethylene. In such cases the vehicle remains in the synthetic thermoplastic polymer after processing. Additionally, the use of an organophilic cation modified clay dispersed in a non-volatile vehicle which may be accompanied by the addition of a volatile liquid organic vehicle which may be substantially completely removed prior to or during processing of the polymer. For best results the modified clay is treated with an organic polar liquid prior to its incorporation in the polymer. Methanol has been found to be suitable for this purpose, but other polar liquids, such as acetone, methyl ethyl ketone and propylene carbonate, can also be used.

Mixtures of the modified clay and one or more flame retardants, particularly such mixture as are obtained by mixing the modified clay and flame retardant(s) in the presence of a volatile organic liquid, adding a polar organic liquid to the mixture, continuing mixing 105 and then evaporating the organic liquids to obtain a mixture of the modified clay and flame retardant(s) in gel form, represent a preferred form of additive composition for the synthetic thermoplastic polymers.

In order that the invention may be more fully understood, the following examples are given by way of illustration only. In these examples all parts are parts by weight per hundred parts by weight of polymer.

Examples 1—16.

A series of low density polyethylene compositions (the polymer being that sold by I.C.I. Ltd. under the trade mark "Alkathene" XDG33) were prepared, the compositions con- 120 taining various amounts of organophilic cation modified clays and flame retardants and the flammability and "dripping rate" of the compositions and of the polymer without these additives were determined.

The compositions were prepared as follows. The modified clay and the flame retardant(s) were added to toluene and the mixture was stirred with a high speed stirrer for 5 minutes,

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methanol was then added and the mixture stirred for a further 5 minutes prior to colloid milling in a mill having a clearance of 0.002 inch. The resultant gel was stored in an oven at 90° C for 12 hours to remove the toluene and methanol.

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The low density polyethylene was added to a mill and the previously prepared gel was added after the polymer had fluxed. The crepe was cross blended frequently and mixing was continued for 5 minutes. The crepe was cut off from the mill and pressed into sheets of inch thickness. The mill temperature was 155° C and the pressing temperature was 145° C.

The test for flammability was essentially as described in A.S.T.M. D635—56T with the following modifications:—

(i) Wire gauze was not positioned below

20 the sample during testing.

(ii) 3 specimens having a length of 4½ in.,

a width of 1 in and a thickness of 1 in (cut

a width of $\frac{1}{2}$ in. and a thickness of $\frac{1}{2}$ in. (cut from the $\frac{1}{2}$ in. thick sheets mentioned above) of each composition were tested. Each specimen was marked by scribing three lines, respectively $\frac{1}{2}$ in., $2\frac{1}{2}$ in. and $4\frac{1}{2}$ in. from one end, thereon.

(iii) The free end of each specimen was ignited with the burner flame and the stop-clock was started when the bulk of the flame reached the first mark, \(\frac{1}{4}\) in. from the free end. The time, in seconds, until the bulk of the flame reached the third mark, $4\frac{1}{4}$ in. from the free end, was measured.

(iv) When the flame reached the second 35 mark, 2½ in. from the free end, the number of drips that fell in 15 seconds was counted and this was recorded as the dripping rate.

(In Examples 13—16 inclusive, the specimens had a length of 6 in., a width of $\frac{1}{3}$ in. 40 and a thickness of 0.006 in., and each specimen was marked by scribing two lines, one at 1 in. and the other at 5 in. from one end of the specimen. The dripping rate was taken 75 seconds after the flame had reached the first 45 mark).

(v) If the specimen was self-extinguishing, in order to obtain a dripping rate, the burner flame was applied so as to keep the specimen burning; (those examples in which this was 50 necessary are marked with an asterisk).

(vi) Testing was discontinued after 3 specimens had been tested and found to be within the following limits:—

for dripping rate $0-5 \pm 1$ 55 $5-10 \pm 2$ $10-20 \pm 3$ $20-40 \pm 4$ for burning rate ± 10 seconds

The average of the burning rate and the dripping rate for the 3 specimens is recorded in the following table; in those cases where the interval between successive drips was greater than 15 seconds, the dripping rate is given as

The results obtained were as follows:—

	Dripping Rate	continuous	12		12		8		9		-	⊽	マ	7	10	
Rate of	Burning (secs.)	220	177		190		360		225		177	185	175	170	220	
	Ratio I:II	ı	1:7.5		1:5		1:7.5		1:5		1:2.5	1:1.25	1:2.5	1:1.25	1:2.5	
Retardant	Parts	1	10	5	5	ν ·	10	ស	5	٧n	አ ባ	2.5	5	2.5	2	
II Flame	Type	1	"Cereclor" P50	Sb _B O ₈	"Cereclor"	Sb ₂ O ₃	"Cerecior"	Space	"Cerector"	Sb ₂ O ₃	T.C.E.P.	T.C.E.P.	T.T.P.	T.T.P.	NH,Br	
Organophilic cation modified clay	Parts	J	2		2		2		2		2	2	2	2	2	-1
I Organoph modified	Type	-	B.38		B.38		B.38		B.38		B.38	B.38	B.38	B.38	B.34	
Fxamule	No.	Blank 1			7		*		4*		ۍ	9	7	8	6	

Ç	Rate	7	⊽	10	11	4	4	7
Rate of	(secs)	135	160	210	150	150	105	65
	Ratio I:II	1:2	1:2	1:1	1:1	1:1	I: I	1:1
Retardant	Parts	4	8	7	0.5	1	2.5	5
II Flame	Type	T.C.E.P.	T.C.E.P.	"Cerector" P50	"Cerector" P50	"Cerector" P50	"Cereclor" P50	"Cereclor" P50
I Organophilic cation modified clay	Parts	2	ኞ	4	0.5	1	2.5	5
I Organophilic c modified clay	Type	B.Sp	B.Sp	'B.Sp	B.38	B.38	B.38	B.38
Roomelle	No.	10	11	12	13	14	15	16

In the foregoing and the following tables:---

4 = dimethyldioctadecyl ammonium montmorillonite

B38 = dimethyldioctadecyl amnonium hectorite

BSp = dimethyldioctadecyl ammonium sepiolite

"Cerecior" = halogenated hydrocarbon

T.C.E.P. = tris-β-chloroethyl phosphate

T.T.P. = tritolyl phosphate

Examples 17 and 18.

Two high density polyethylene compositionsflammability and dripping rate of the compositions that solid under the tradetions and of the polymer without the additives mark "Rigidex") containing different amountswere determined as described for Examples of a modified clay and a flame retardant were1—12, prepared as described for Examples 1—16, The results obtained were as follows:—except that the mill temperature was 170° C

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Drinning	Rate	continuous	14	П
Rate of	(ses)	360	300	285
	Ratio I:II	1	1:1	1:1
Retardant	Parts	•	7	4
II Flame	Type	.1	"Cerector" 70	"Cereclor" 70
ulic cation clay	Parts		2	4
I Organophilic cation modified clay	Type	[B.Sp	B.Sp
o Cumon	No.	Blank 2	LT	18

25 perature was 170° C, and the flammability and dripping rate of the compositions and of the polymer without the additives were determined as described for Examples

EXAMPLES 19-23

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were as follows: The results obtained A series of polypropylene compositions containing modified clay and various flame retardants were prepared as described for Examples 1—16, except that the mill temperature was 170° C and the pressing tem-

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Drípoine	Rate	continuous	20	2-3	. 1—2	√	1
Rate of	(ses)	264	220	230	205	150	170
	Кайо І:П	1	1:5	1:7.5	1:5	1:2.5	1:1.25
Retardant	Parts	1	10	10	רא רא	5	2.5
II Flame	Type	j	"Cereclor" · 70	"Cereclor" 70 Sb ₀ O ₃	"Cereclor" 70 Sb ₂ O ₃	T.C.E.P.	T.C.E.P.
I Organophilic cation modified clay	Parts	i de la companya de l	2	2		.2	2
I , Organophilic ; modified clay	Type	1.	B.38	B.38	B.38	B.38	B.38
Ç	No.	Blank 3	61	20	21	22	23

8 of the composition containing an organophilic cation modified zeolite clay as the modified clay. The polymer was fluxed in an internal mixer at 240° C for 2 minutes, the pre-pre-pared modified clay/flame retardant gel was then added and mixing was continued for a further 5 minutes. The resultant mass was removed and pressed at 230° C into sheets \$\frac{1}{2}\$ in. were cut from these sheets for testing.

The results obtained were as follows:—

Example 24 and 25.

Two nylon 11 compositions containing modified clay and flame retardant were prethese compositions and of the unmodified polymer were determined as described for Examples 1—12.

To prepare the compositions, a gel of the modified clay and the flame retardant was prepared as described for Examples 1—16, execept that methanol was omitted in the case

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	Dripping Rate	Continuous	7	5
Rate of	Burning (secs)	173	206	163
	Ratio I:II	ı	1:1,25	1:1.25
Retardant	Parts	1	6	6
II Flame	Type	ı	T.C.B.P.	T.C.B.P.
nilic cation d clay	Parts	1	7	4
I Organophilic cation modified clay	Type	I	B.38	B.27
Example	No.	Blank 4	24	25

B.27 = an organophilic zeolitic clay.

Example 26.

100 Parts of a toughened polystyrene (sold the Registered Trade Mark "Carinex" polystyrene (sold the natural N 3001 TGX by the Shell Chemical do. Ltd.,) was milled on a heated two roll in mill at 160° C until fluxed. The modified clay the and fire retardant were added gradually and

mixing was continued for 5 minutes after all the additive had been incorporated into the polymer. The rough crepe of polymer thus produced was pressed into sheets 6 in. × 4 in.

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<u></u>	•••	-		
Dripping	Rate	20	.8	8
	Kago 1:11	1	İ	1.2.5
II. Flame Retardant	Parts	1	เก	5
II. Flame	Type]	TCEP	TCEP
I. Organophilic cation modified clay	Parts		ı	2
I. Organop modifie	Type] .	1.	B.38
Example No.	Example No.		Blank 6	26

WHAT WE CLAIM IS:—

1. A polymer composition comprising a polyamide, polystyrene or a polyolefin derived from a straight chain olefin and, incorporated therein, one or more flame retardants and up to 10%, based on the weight of the polymer, of an organophilic cation modified clay.

2. A polymer composition according to claim 1, which comprises a total of from 0.5 to 50%, based on the weight of the polymer,

of flame retardant.

3. A polymer composition according to claim 1 or 2, in which the flame retardant is a chlorinated parassin, tris-β-chloroethyl phosphate, tribromophenol, ammonium chloride, ammonium bromide, a phosphoric acid ester, antimony trioxide or arsenic trioxide.

4. A polymer composition according to any of claims 1 to 3, which comprises from 0.5 to 5%, based on the weight of the polymer,

of organophilic cation modified clay.

5. A polymer composition according to any of claims 1 to 4, in which the organophilic cation modified clay is dimethyl dioctadecyl ammonium sepiolite, octadecyl ammonium montmorillonite, dimethyl dioctadecyl ammonium montmorillonite, or dimethyl dioctadecyl ammonium hectorite.

6. A polymer composition according to any of claims 1 to 4, in which the organophilic cation modified clay is a cationic clay of high cation exchange capacity which is associated with an aliphatic amine containing at least 12 carbon atoms or a derivative thereof.

7. A polymer composition according to claim I substantially as herein described in any of Examples 1 to 25.

8. A polymer composition according to claim 1 substantially as herein described in Example

40 26.

9. An additive composition for imparting

flame-retardancy and drip-resistant properties to synthetic thermoplastic polymers as herein defined, which comprises an admixture of one or more flame retardants and an organophilic cation modified clay, said mixture containing a total of from 1 to 10 parts by weight of flame retardant per part by weight of modified clay.

10. An additive composition according to claim 9, in which the flame retardant is a chlorinated parattin, tris-\beta-chloroethy! phosphate, tribromophenol, ammonium chloride, ammonium bromide, a phosphoric acid ester, antimony trioxide or arsenic trioxide.

11. An additive composition according to claim 9 or 10, in which the organophilic cation modified clay is dimethyl dioctadecyl ammonium sepiolite, octadecyl ammonium montmorillonite, dimethyl dioctadecyl ammonium montmorillonite, or dimethyl dioctadecyl ammonium hectorite.

12. An additive composition according to claim 9 or 10, in which the organophilic cation modified clay is a cationic clay of high cation exchange capacity which is associated with an 65 aliphatic amine containing at least 12 carbon

atoms or a derivative thereof.

13. An additive composition according to any of claims 9 to 11 which has been prepared by mixing the modified clay and flame retard- 70 ant(s) in the presence of a volatile organic liquid, adding a polar organic liquid to the mixture, continuing mixing and then evaporating the organic liquids to obtain the mixture in gel form.

> A. A. THORNTON & CO., Chartered Patent Agents, Northumberland House, 303—306, High Holborn, London, W.C.1. For the Applicants.

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